

MIA-002: A randomised, vaginal microbicide trial assessing the safety of PRO 2000/5 gel (P) versus vehicle placebo in Uganda

Submission date
11/08/2005

Recruitment status
No longer recruiting

☐ Prospectively registered

☐ Protocol

Registration date
15/09/2005

Overall study status
Completed

☐ Statistical analysis plan

☒ Results

Last Edited
25/05/2010

Condition category
Infections and Infestations

☐ Individual participant data

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

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Additional identifiers

Protocol serial number

MIA-002 Version 3

Study information

Scientific Title

Acronym

MIA: Microbicides Initiative in Africa

Study objectives

That PRO 2000/5 (P) gel in 0.5% and 2% formulations are as safe and acceptable for women to use as a vehicle placebo gel

Ethics approval required

Old ethics approval format

Ethics approval(s)

Not provided at time of registration

Study design

Randomised controlled trial

Primary study design

Interventional

Study type(s)

Prevention

Health condition(s) or problem(s) studied

HIV-1

Interventions

Two active study products (0.5% and 2% PRO 2000/5 Gel [P]) and a matched vehicle placebo (Placebo Gel [P]) inserted intra-vaginally twice a day for 28 days

Intervention Type

Drug

Phase

Phase II

Drug/device/biological/vaccine name(s)

PRO 2000/5 gel

Primary outcome(s)

The primary end-points are local and systemic safety parameters, namely:

1. Deep (any size) or extensive superficial (greater than or equal to 4 times the size of the tip of a 5 x 10 mm cotton-tipped swab) genital epithelial disruption visible on naked eye examination or colposcopy
2. The appearance of a coagulation abnormality which is considered clinically relevant by the local investigator/Trial Management Group

Key secondary outcome(s)

The secondary end-points are:

1. Grade 3 clinical or laboratory adverse event confirmed on examination or repeat testing respectively, thought to be possibly or probably related to gel

2. Grade 3 unexpected vaginal bleeding as reported at interview or on a diary card or recorded on examination
3. Grade 1 unexpected vaginal bleeding not due to menses
4. Acceptability of gel as assessed by a semi-structured questionnaire
5. Alterations in vaginal flora assessed by Nugent score performed on Gram-stained slides

Completion date

17/11/2004

Eligibility

Key inclusion criteria

1. Healthy* women aged between 18 and 45
2. Sexually active and likely to remain so for the duration of the study at a minimum rate of twice per week
3. Willing to undergo a genital infection screen
4. Willing to undergo a human immunodeficiency virus (HIV) test**
5. Willing to accept health education about condoms and to be supplied with condoms to be used at every episode of sexual intercourse during the study, and has used condoms before
6. Able to give informed consent
7. Either HIV negative or HIV positive in a monogamous sexual relationship with another person who is also HIV positive and who will give his signed consent to say he has been informed and understands about the trial

* HIV-seropositive women will be eligible providing they fulfil all other criteria including exclusion 10, and have a primary partner who is also HIV-seropositive. Antiretroviral therapy is permitted provided it has been stable for 2 months prior to enrolment and is not expected to change during participation in the study

** Unnecessary if HIV-positivity documented in medical records at Nsambya Hospital

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Upper age limit

45 years

Sex

Female

Key exclusion criteria

1. Pregnant, wanting to become pregnant or within 6 weeks postpartum
2. Current genital tract epithelial ulceration/disruption
3. Untreated gonococcal, chlamydial, or trichomonal infection, syphilis or symptomatic bacterial vaginosis
4. Is HIV positive and in a sexual relationship with someone who is not HIV infected, who will not have an HIV test before the trial, and/or who will not sign a consent form for the trial
5. Abnormal (grade II) haematology, biochemistry
6. Cervical intraepithelial neoplasia (CIN) greater than or equal to CIN II within 3 months
7. Acute/subacute pelvic inflammatory disease
8. Clinical coagulation disorder
9. Latex allergy
10. Current, recent (within 2 weeks) or on-going ill health that necessitates drug treatment (other than prophylaxis) or attendance at hospital***
11. Post-coital or intermenstrual bleeding in the past 3 months
12. (If post-natal) Persistent abnormal vaginal discharge
13. No reported use of condoms between screening and enrolment
14. Having participated in another microbicide trial in previous 30 days
15. Considered unlikely to be able to comply with protocol

*** Except for HIV positive women from the HIV clinic, where the same exclusion applies without the phrase 'or ongoing'; women may enter the study following the initiation of appropriate treatment

Date of first enrolment

11/06/2003

Date of final enrolment

17/11/2004

Locations

Countries of recruitment

United Kingdom

England

Uganda

Study participating centre

222 Euston Road

London

United Kingdom

NW1 2DA

Sponsor information

Organisation

Imperial College London (UK)

ROR

<https://ror.org/041kmwe10>

Funder(s)**Funder type**

Government

Funder Name

The project was funded by the European Commission

Results and Publications**Individual participant data (IPD) sharing plan****IPD sharing plan summary**

Not provided at time of registration

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	01/06/2010		Yes	No